Application No.: 09/117,838 Amendment dated: August 1, 2008 Reply to Office Action of April 1, 2008 Attorney Docket No.: 0075.0006US1

Amendments to the Specification

Please, replace the last paragraph on page 3 with the following paragraph:

-- Conceptually, the present invention claims a novel category (class) of medicinal preparations and/or medicinal forms that can be specified as "Bipathic", combining therapeutic values of medicinal substance in therapeutic dose and potentiated homeopathic preparation chemically homogeneous (in original formula or composition of the initial substance) but different in mechanism of action on the organism. This combination promotes biological activation and induces positive morphological and functional changes in form of "systemic adaptation" responsible for increased therapeutic efficiency of the active medicinal substance with reduced risk of patients' individual reactions and undesirable adverse after-effects.—

Please, replace the paragraph of Example 1 with the following paragraph:

-- Prior to transfer of bioenergetic information, 10 ml of 0.5% solution of atropine sulphate (medicine in therapeutic dose) as a carrier, and as a bioactive substance, potentiated preparation Atropini Sulfati C30 obtained by multiplesuccessive dilution and shaking in accordance with homeopathic method, are placed in two separate containers mounted on current-conducting plates connected via a circuit of a known recorder of information signal. During bioenergetic information exchange, information about on homeopathically potentiated initial active substance - atropine - is transferred to the carrier. Potentiated atropine The initial active substance has chemical formula identical to that of the carrier and possesses field withcertain frequency spectrum. The obtained medicine is applied in ophthalmology as a mydriatic for diagnosis and treatment of inflammatory conditions; it is devoid of accommodation paralysis as an adverse effect. –

Change(s) applied to document,

/G.D./

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Please, replace the paragraph of Example I with the following paragraph:

-- 0.01 g of potentiated homeopathic preparation Acidum Salicylicum is pressed into a pill containing 0.5 g of acetylsalicylic acid. The former is produced in accordance with homeopathic method by saturating a neutral substance, lactose, with solution of Acidum Salicylicum in C30 potency. By potentiating, the initial substance - acetylsalicylic acid - is bioenergetically transformed in accordance with homeopathic